Amendments to the Specification:

Please delete the Abstract of the Disclosure on page 1, from lines 8-20 of the specification and insert the following paragraph on a new page 105 following the claims:

-- ABSTRACT OF THE DISCLOSURE

This invention relates to newly identified polynucleotides and polypeptides, and their production and uses, as well as their variants, agonists and antagonists, and their uses. In particular, the invention relates to polynucleotides and polypeptides of a *Staphylococcus aureus* (S. aureus) DnaI related protein, as well as its variants, hereinafter referred to as "S. aureus DnaI", "S. aureus DnaI polypeptide(s)", and "S. aureus dnaI polynucleotides" as the case may be. The invention also relates to a specific interaction between the S. aureus DnaI related protein or specific regions thereof, and a growth-inhibitory protein encoded by the S. aureus bacteriophage 77 genome. The phage open reading frame (ORF) product interacts with amino acids 150 – 313 of S. aureus DnaI polypeptide, and the invention relates to the use of this interaction target site as the basis of drug screening assays. Accordingly, the invention provides a method for the inhibition of bacterial growth, and the treatment of bacterial infection via the inhibition of DnaI. --

Please replace the paragraph on page 11, lines 12-13, with the following description.

--Fig. 2<u>A-2N</u> shows the complete nucleotide sequence of the *S. aureus* bacteriophage 77 genome (SEQ ID NO: 3).--

Please replace the paragraph on page 12, lines 1-7 with the following paragraph:

--Fig. 6A-E shows sequences and alignments of B. subtilis DnaC sequences with the homologous sequences from S. aureus. 6C) shows an alignment of B. subtilis dnaC polynucleotide sequence (SEQ ID NO: 6) with the homologous S. aureus dnaC polynucleotide sequence (SEQ ID NO: 7) identified by BLAST searching the S. aureus database at http://www.tigr.org The Institute of Genomic Research (TIGR) web site with the B. subtilis dnaC sequence. 6E) shows an alignment of B. subtilis DnaC amino acid sequence (SEQ. ID NO: 8) with the predicted amino acid sequence of the polypeptide (SEQ ID NO: 9) encoded by the S. aureus dnaC polynucleotide sequence shown in Fig. 6B.--

Please replace the partial paragraph on page 45, lines 1-8 with the following paragraph:

--http://www.genome.ou.edu/staph.html The University of Oklahoma's Advanced Center for Genome Technology web site. One sequence contig of 4850 nucleotides in length (Contig 981), when converted into amino acid sequence, contained within it the similar amino acid sequence GHVPELYVDNNR (SEQ ID NO: 11; Fig. 5). This tentative identification of the candidate

protein was then confirmed upon *in silico* tryptic digestion of the open reading frame found in the contig (Fig. 5). The obtained PSD/CID spectra for tryptic peptides with monoisotopic MH+ masses of 1351.8, 1412.7, and 1617.8 Da were similar to the predicted PSD/CID fragmentation patterns of the tryptic peptides with monoisotopic MH+ masses of 1351.8 and 1617.8 Da found in the contig's +3 open reading frame (Fig. 5).

Please replace the paragraph on page 46, lines 18-23 with the following paragraph:

--Databases were searched for *S. aureus* genes which may be related to the *B. subtilis* dnaC gene. Utilizing the *B. subtilis* amino acid sequence for DnaC (Accession Number P37469), a BLAST search was performed of the Staphylococcus database at http://www.tigr.org The Institute of Genomic Research (TIGR) web site and revealed the presence of an ORF within the *S. aureus* genome encoding a related protein. The nucleotide sequence and corresponding protein sequence are presented in Fig. 6AB (SEQ ID NO: 7) and Fig. 6BD (SEQ ID NO: 9), respectively.--

Please replace the partial paragraph on page 84, lines 1-5 with the following partial paragraph:

--genome were sequenced at least once from both directions on two separate clones. Sequence contigs were assembled using Sequencher 3.1 software (GeneCodes) (Fig. 2). An implementation of the publicly available program SEQUIN, available for download at http://negi.nlm.nih.gov/sequin/ The United States National Library of Medicine's web site, was used on phage genome sequence to identify all putative ORFs larger than 33 codons (Fig. 3). --

Please replace Table 1 on the page immediately following the claims, with the following Table 1.

--Table 1 Similarities in sequence between the DnaI homolog and sequences deposited in public database.

Sequences producing significant alignments:

	core	E
	(bits)	Value
gi 140025 sp P06567 DNAI_BACSU PRIMOSOMAL PROTEIN DNAI >gi 2797	231	5e-60
gi 468268 (M15183) ORFY [Bacillus subtilis]	125	5e-28
gi 2072367 emb CAA70453 (Y09255) primosomal protein DnaI [Baci	84	1e-15
gi 530419 emb CAA83732 (Z33058)GTP bind. CD48/PAS1/ SEC18 fam	67	2e-10
gi 2983431 (AE000713) DNA replication protein DnaC [Aquifex aeo	52	4e-06
gi 1176732 sp P45910 YQAM_BACSU HYPOTHETICAL 36.1 KD PROTEIN IN	50	2e-05
gi 2127076 pir I40411 hypothetical protein 5 (xre region) - Ba	48	1e-04
gi 1722861 sp P39782 XKDC_BACSU PHAGE-LIKE ELEMENT PBSX PROTEIN	48	1e-04
gi 1353529 (U38906) ORF12 [Bacteriophage rlt]	46	4e-04
gi 2983000 (AE000683) chromosome replication initiator protein	45	8e-04

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>qi|140025|sp|P06567|DNAI_BACSU PRIMOSOMAL PROTEIN DNAI
           >qi|279708|pir||IQBS44 dnaA protein homolog, 44K -
           Bacillus subtilis >gi|39881|emb|CAA28633| (X04963) ORF
           311 (AA 1-311) [Bacillus subtilis]
           >gi|1769996|emb|CAA99605| (Z75208) replication protein
           [Bacillus subtilis] >gi|2293281 (AF008220) DnaI
           [Bacillus subtilis] >gi|2635363|emb|CAB14858| (Z99118)
           helicase loader [Bacillus subtilis]
           Length = 311
Score = 231 bits (583), Expect = 5e-60
Identities = 120/280 (42%), Positives = 177/280 (62%), Gaps = 2/280 (0%)
Query: 35 DPDVKQFLEAHRAELTNAMIDEDLNVLQEYKDQQKHYDG-HKFADCPNFVKGHVPELYVD 93
          Sbjct: 31 DQDVQAFLKENEEVIDQKMIEKSLNKLYEYIEQSKNCSYCSEDENCNNLLEGYHPKLVVN 90
Query: 94 NNRIKIRYLQCPCKIKYDEERFEAELITSHHMQRDTLNAKLKDIYMNHRDRLDVAMAADD 153
             I I Y +CP K K D+++ + L+ S ++Q+D L A + + ++ RL +
Sbjct: 91 GRSIDIEYYECPVKRKLDQQKKQQSLMKSMYIQQDLLGATFQQVDISDPSRLAMFQHVTD 150
Query: 154 ICTAITNGEQVKGLYLYGPFGTGKSFILGAIANQLKSKKVRSTIIYLPEFIRTLKGGFKD 213
                   + KGLYLYG FG GK+F+L AIAN+L K+ S I+Y+PEF+R LK
Sbjct: 151 FLKSYNETGKGKGLYLYGKFGVGKTFMLAAIANELAEKEYSSMIVYVPEFVRELKNSLQD 210
Query: 214 GSFEKKLHRVREANILMLDDIGAEEVTPWVRDEVIGPLLHYRMVHELPTFFSSNFDYSEL 273
           + E+KL+ V+ +LMLDDIGAE +T WVRDEVIG +L +RM +LPTFFSSNF EL
Sbjct: 211 QTLEEKLNMVKTTPVLMLDDIGAESMTSWVRDEVIGTVLQHRMSQQLPTFFSSNFSPDEL 270
Query: 274 EHHLAMTRDGE-EKTKAARIIERVKSLSTPYFLSGENFRN 312
               ++ GE E+ KAAR++ER+ L+ P L GEN R+
          +HH
Sbjct: 271 KHHFTYSQRGEKEEVKAARLMERILYLAAPIRLDGENRRH 310
> gi | 468268 (M15183) ORFY [Bacillus subtilis]
           Length = 207
 Score = 125 bits (310), Expect = 5e-28
 Identities = 67/190 (35%), Positives = 105/190 (55%), Gaps = 1/190 (0%)
Query: 20 DFXXXXXXXXXXXXXNDPDVKQFLEAHRAELTNAMIDEDLNVLQEYKDQQKHYDG-HKFAD 78
                        D DV+ FL+ + MI++ LN L EY +Q K+
          DF
         DFQNRLEQTKEKVMKDQDVQAFLKENEEVIDQKMIEKSLNKLYEYIEQSKNCSYCSEDEN 75
Sbjct: 16
Query: 79 CPNFVKGHVPELYVDNNRIKIRYLQCPCKIKYDEERFEAELITSHHMQRDTLNAKLKDIY 138
          C N ++G+ P+L V+ I I Y +CP K K D+++ + L+ S ++Q+D L A + +
Sbjct: 76 CNNLLEGYHPKLVVNGRSIDIEYYECPVKRKLDQQKKQQSLMKSMYIQQDLLGATFQQVD 135
Query: 139 MNHRDRLDVAMAADDICTAITNGEQVKGLYLYGPFGTGKSFILGAIANQLKSKKVRSTII 198
              RL + D + + KGLYLYG FG GK+F+L AIAN+L K+ S I+
Sbjct: 136 ISDPSRLAMFQHVTDFLKSYNETGKGKGLYLYGKFGVGKTFMLAAIANELAEKEYSSMIV 195
Query: 199 YLPEFIRTLK 208
          Y+PEF+R LK
Sbjct: 196 YVPEFVRELK 205--
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